
BioMax Environmental

Environmental Consulting and Industrial Hygiene Services

April 3rd, 2008

Mr. Doug Button
Deputy Director
Real Estate Services Division
707 Third Street - 8th Floor
West Sacramento, CA 95605

Post Mitigation Assessment
Department of General Services Board of Equalization Building
24th Floor Break Room Areas
450 N. Street
Sacramento, California

BioMax Environmental, LLC (BioMax) is pleased to provide The Department of General Services (DGS) with this letter summary report detailing BioMax's findings and recommendations pertaining to our post mitigation microbial inspection and sampling assessment services provided within the 24th Floor Break room areas present within the Board of Equalization (BOE) building (subject building) located at 450 N Street, Sacramento, California. BioMax understands that these post mitigation microbial inspection and clearance sampling assessment services were contracted with BioMax, at your request, in an effort to review and verify the successful completion of microbial mitigative efforts performed by your restoration contractor, JLS Environmental, Inc., within the previously identified mold damaged and moisture impacted areas within the 24th floor break room areas of the subject building. These microbial clearance assessment services were requested following the discovery of moisture and mold damage materials visually identified within the noted break rooms and summarized within BioMax's summary report entitled Mitigation Procedures for Moisture Impacted Break Room Areas, dated March 19th, 2008.

Hence, these post mitigation assessment services are intended to assess the current site conditions wherein mitigative activities were performed to address prior moisture related damages and impacts. Following the completion of prescribed mitigative activities performed by your selected contractor, JLS, Mr. Michael A. Polkalla, CIH, REA of BioMax performed a post mitigation site inspection and sampling assessment within the affected areas of the subject building areas as noted below. BioMax's findings and conclusions pertaining to our post mitigation sampling assessment are summarized herein.

These post mitigation microbial clearance assessment services, thereby, are intended to provide a professional evaluation supported by technical sampling information verifying physical

conditions wherein the successful completion of microbial removal and decontamination within the affected areas has been achieved.

SITE OBSERVATIONS

Site inspection and post mitigation assessment sampling activities were performed on Wednesday, March 26th, 2008 wherein site access into contained and non-contained areas was facilitated by Mr. Rick Boggs of JLS. On this day, Mr. Michael A. Polkabila, CIH, REA of BioMax performed a visual site inspection of each of the containment system barriers associated with the two 24th floor break rooms (Rooms 2424 and 2402) and collected a series of airborne samples within and surrounding the areas of concern the subject structures as noted below.

On-site inspection and clearance sampling assessment activities were performed by Mr. Michael A. Polkabila, CIH, REA, of BioMax in accordance with currently recognized microbial assessment and sampling guideline procedures. Mr. Polkabila has been certified in the Comprehensive Practice of Industrial Hygiene by the American Board of Industrial Hygiene and holds the right to the designation "Certified Industrial Hygienist" (CIH) under certification number CP 7104. Mr. Polkabila is also certified by the California Environmental Protection Agency (Cal/EPA) as a Class I Registered Environmental Assessor (REA) under Cal/EPA certification number 05011. A summary of significant notations and observations gathered during BioMax's site inspection and clearance assessment within the subject facility are compiled as follows:

1. At the time of our site inspection and clearance sampling assessment performed on February 19th, 2008 ambient outdoor conditions both prior to and following our interior assessment consisted of sunny and mild conditions with an outdoor temperatures range between 46 and 66 degrees F and relative humidity range between 34 and 55 %. Predominant winds were noted at approximately 0-5 knots from the northwesterly direction at the time of our assessment. Interior environmental conditions within the sampled 24th Floor areas consisted of a temperature range between 74 and 78 degrees F with relative humidity range of 25 to 28 percent.
2. Interior containments whereby microbial mitigative activities were previously performed by JLS included doorway as well as ceiling plastic barriers erected within the impacted break room areas including Room 2424 and 2402. Based on our inspection and review of observations within and surrounding each of the containment areas, BioMax concluded that such barrier systems provided evidence of appropriate control barriers and site protections at the time of our post mitigation assessment. A detailed site map indicating the delineation of established (as-constructed) containment systems utilized during this procedure may be referenced from JLS site records, as necessary.
3. Based on our post mitigation inspection within and surrounding the containment areas noted above, BioMax documented the absence of visible interior indications of elevated residual moisture and/or microbial indicators (such as staining, delamination, etc.) within the

remaining exposed interior walls, wall framing, and wall cavities following the performance of material removal and mitigative measures.

4. Utilization of a TraMex hand-held inductive moisture meter indicated normal moisture content within all remaining walls and building materials inspected within each of the sampled containment break room areas at the time of our assessment.
5. At the time of our post mitigation inspection, containment system encompassing each of the interior affected break room areas were observed and verified under appropriate posting and negative pressure differential. Worker and equipment entry and exit chambers comprised of a series of zippered plastic access doorways were also observed attached to the noted containment break room barrier consistent with BioMax's written mitigation recommendation protocols.
6. As prescribed, all identified affected interior cabinetry and wallboard building materials had been removed from each of the affected areas exposing interior wall cavity framing (metal) as well as underlayment wallboard siding materials. Upon post mitigation inspection, all remaining exposed building materials associated with the interior wall systems exhibited no significant staining and/or elevated mold growth following the completion of prescribed physical mold removal and chemical decontamination procedures performed by the selected mitigation contractor on the surfaces of such exposed building materials.
7. In conjunction with our visual inspection, BioMax collected series airborne samples within and outside each of the containment areas noted in Table 1 below for subsequent comparative analysis. Such samples collected within the interior containment area were performed in an effort to identify and quantify the presence of any potential significant fugitive airborne mold spores present within (and surrounding) the containment systems following the mitigative effort.
8. BioMax also collected a series of digital images during this post mitigative inspection and sampling assessment activities to document the conditions and significant site observations gathered at this time. Such images are provided as an attachment to this summary report for further reference, as necessary.

SAMPLING PROCEDURES

On-site inspection and sampling assessment activities were conducted by Mr. Michael A. Polkaba, CIH, REA, of BioMax Environmental on March 26th, 2008. All sampling equipment, supplies, calibration materials, and collection media were provided by BioMax as part of the performance of this scope of work. Sample collection procedures and methods were performed using aseptic sampling methods following techniques prescribed by the contracted analytical laboratory.

Spore Trap Airborne Microbial and Particulate Sampling:

The collection of airborne Spore Trap microbial samples was achieved using Zefon Air-O-Cell sampling cassette collection devices placed in each of the areas identified in Table 1. A total of ten (10) airborne Spore Trap samples were collected within and outside the containment areas at a height of approximately four feet above ground level using a tripod mounted Quick Take 15 air sampling pump manufactured by SKC. Samples were collected at a calibrated flow rate of 15 liters per minute for a total of five minutes per sample. Resultant total sample volumes, therefore, corresponded to 75 liters collected for each sample. Field calibration of the SKC air sampling pump was conducted and recorded prior to sampling activities using a field rotometer calibrated with a Bios Drycal primary standard flow meter. All spore trap air sampling and analytical procedures were performed in accordance with prescribed manufacturer guidelines as well as applicable professional certified industrial hygiene indoor air quality microbial investigation procedures and certified industrial hygiene practices.

Additional exterior samples were also similarly collected and analyzed as part of the collection of interior samples in an effort to identify and quantify normal background microbial taxa (types), rank order, and corresponding airborne spore levels present at the time of this assessment. It should be noted that all Heating Ventilation and Air Conditioning (HVAC) systems on the 24th floor had been deactivated in October of 2007. Hence, the collection of airborne samples performed during this clearance assessment were representative of building conditions present within each of the subject areas sampled at the time of this assessment. Sampling collection activities performed during this study included the collection of identifiable airborne microbial contaminants within the representative areas noted in Table 1 below:

Table 1. Airborne Spore Trap Sampling Locations:

Air Sample Number	Spore Trap Air Sampling Location
13430732	Ambient outside location (Main Entry Level)
13430735	Ambient outside sample from 12 th Floor SE balcony
13430710	Ambient 23 rd Floor Western Balcony
13430739	Hallway outside containment near 2424
13430694	Break Room Containment 2424 (L)
13430729	Break Room Containment 2424 (R)
13430705	Hallway near 2402 (outside containment)
13430701	Break Room Containment 2402 (L)
13430718	Break Room Containment 402 (R)

Air Sample Number	Spore Trap Air Sampling Location
13430706	Ambient 23 rd Floor Western Balcony

At the conclusion of sampling activities, preparation and shipping of the collected samples were accomplished in accordance with standard industrial hygiene chain of custody (COC) documentation procedures and quality assurance/quality control practices. Once collected, labeled, and recorded, all samples were double sealed within airtight plastic Ziploc shipping containers and transported via Federal Express Priority Mail to Environmental Microbial Laboratories (EMLabs) in San Bruno, California. EMLabs holds current applicable analytical accreditation and specializes in microbial analytical procedures. Sampling and chain of custody records are provided as an attachment to this letter report for further reference.

ANALYTICAL FINDINGS AND CONCLUSIONS

Airborne Spore Trap Findings:

Laboratory analytical methods for the identification and enumeration of microbial (mold) taxa and particulate contaminants were conducted in accordance with prescribed analytical procedures and quality control/assurance measures. Original laboratory results including the enumeration of recognizable microbial spore and particulate types are also attached to this letter report for further detail. Analytical comments provided by the microbial laboratory regarding relative levels are noted as a semi-quantitative assessment based on historical and regional data. A summary of airborne Spore Trap microbial (mold) and particulate findings within and surrounding each of the subject Break room areas are presented in Table 2 below:

Table 2. Summary of Airborne Microbial and Particulate Findings

Location Desc.	Total Mold Spores (Cts/m ³)	Background Debris (scale of 1-4)	Skin Cell Fragments (scale of 1-4)
Ambient outside location (Main Entry Level)	1,281	2+	<1+
Ambient outside sample from 12 th Floor SE balcony	334	2+	<1+
Ambient 23 rd Floor Western Balcony AM	386	2+	<1+
Hallway outside containment near 2424	107	3+	2+

Location Desc.	Total Mold Spores (Cts/m ³)	Background Debris (scale of 1-4)	Stain Cell Fragments (scale of 1-4)
Break Room Containment 2424 (L)	107	3+	2+
Break Room Containment 2424 (R)	66	3+	2+
Hallway near 2402 (outside containment)	66	3+	3+
Break Room Containment 2402 (L)	106	3+	1+
Break Room Containment 402 (R)	107	3+	1+
Ambient 23 rd Floor Western Balcony PM	226	2+	<1+

The analytical findings presented in Table 2 indicate the presence of significantly lower concentrations of microbial (mold) spores measured within each of the interior samples collected within and surrounding the subject Break Room areas when compared to the levels currently measured within the samples collected from the corresponding ambient outside environment. Analytical findings also indicate similar fungal taxa distribution (mold types) and rank order (predominant taxa) of molds identified within the mitigated areas as well as the adjacent hallways and worker occupied area samples present outside the noted containment areas. Particularly worthy of note, was the absence of significantly elevated levels of hydrophilic (moisture loving) airborne mold taxa such as *Penicillium*, *Aspergillus*, and *Stachybotrys* as comparatively summarized in this assessment report.

Although there are currently no regulatory standards or limits pertaining to allowable airborne fungal concentrations (for any mold taxa) present in indoor environments, there is a general consensus among indoor air quality experts that microbial contamination found within "typical healthy" living spaces are generally similar in kind and present at levels which are below those found in the corresponding native outside environment. BioMax believes that the absence of elevated residual moisture, absence of significant visible residual mold and/or staining, and relatively fewer total airborne mold levels with typical taxa and rank order distribution following mitigative clean-up activities are consistent with these generally acceptable conditions. BioMax, therefore, believes that these findings provide reasonable evidence indicating that current microbial clean-up measures have successfully mitigated and contained mold contamination within previously affected break room areas and previously affected materials.

Based on these findings, BioMax believes that the current site conditions present within the mitigated break room areas as well as the corresponding analytical data collected and evaluated, following the performance of the recommended mitigative procedures, meets the mitigative clearance criteria established for these activities as presented in BioMax's Post Mitigation Clearance Assessment Protocols dated February 15th, 2008 as reviewed and approved by BOE and their environmental consultant, Hygientech. Therefore, BioMax believes that achievement of such criteria warrants our determination and recommendation that the previously impacted areas may be considered acceptable for normal reconstruction as there are no current significant data or current evidence to the contrary.

Airborne Particulate Findings:

Analytical findings pertaining to the airborne particulates consisting primarily of debris, pollen, and skin cell fragments identified within the collected air samples within and surrounding the previously impacted areas also provide reasonable evidence indicating that current particulate clean-up and mitigative control measures have successfully controlled and contained particulate debris within the identified containment areas to normal background ranges.

Although, there are similarly no currently applicable regulatory standards pertaining to allowable particulate levels with which to compare, it is BioMax's professional opinion that interior particulate levels should continue to be minimized wherever possible. Therefore, additional (and ongoing) recommendations for optional particulate control measures have been provided at the end of this report for client consideration.

RECOMMENDATIONS

Based on the findings and conclusions presented in this report, BioMax believes that the current airborne microbial levels sampled and analyzed from within the identified 24th Floor Break Room areas (Rooms 2424 and 2402) provides no significant evidence of elevated residual microbial contamination or airborne migration following the completion of prescribed microbial mitigative measures. Hence, based on our direct site observations, measurements, and review of these findings at this time, BioMax believes that the previously affected areas located within the noted Break Room areas may be considered acceptable for general reconstruction following prudent reconstruction practices with the implementation of the noted additional measures discussed below. Therefore, based on these findings, BioMax recommends consideration of the following post-mitigation measures and actions:

1. BioMax believes that current airborne microbial (mold) levels and mold types were identified at levels which are believed to fall within generally acceptable ranges and parameters at present. Hence, BioMax recommends that no further airborne microbial sampling activities are warranted within the specific areas sampled and mitigated at this time. Certainly, due to the knowledge that microbial contamination, by nature, may change over time due to additional moisture intrusion, favorable growth conditions, and changing environments, these

recommendations are subject to revision in the event that such conditions and/or environments arise.

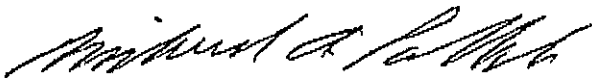
2. During the performance of interior reconstruction activities, BioMax recommends that a qualified and experienced building inspector/contractor be utilized to verify the current functional integrity of all applicable remaining plumbing, flashing, scaling, and drainage systems in accordance with current building codes and construction practices. Any identified deficiencies should be appropriately documented, corrected, and then functionally verified (tested) prior to subsequent reconstruction and commercial use. Certainly, the establishment/installation of any additional engineering controls (as identified through additional professional engineering consultation) should also be performed and implemented in accordance with applicable standards, building codes, and ordinances, as necessary.
3. Due to the anticipated forthcoming renovation and reuse of these areas for break room related activities, as an additional precautionary measure, BioMax recommends that all remaining interior wall cavity wallboard materials be reconstructed utilizing an appropriate grade of sheetrock materials where moisture barrier properties are desired. As a client option, any interior sheetrock surfaces may also receive the application of a spray-on sealant with microbial growth inhibitors prior to wall cavity construction. The application of such sealant (if desired) should certainly be performed by your selected contractor in accordance with all product manufacturer's use specifications and application guidelines.
4. BioMax recommends that all reconstruction of interior structural building materials should only be undertaken utilizing high quality, visibly clean (hand selected) construction grade building materials obtained from reputable commercial sources and which are believed and visually free from elevated microbial contamination and/or elevated moisture content. Building materials, which are notably moist and/or visibly stained, should not be used during the reconstruction undertaken within the subject residence. BioMax recommends that all current plastic barriers (as established during this mitigative effort) may remain during such reconstruction so as to minimize the potential transmission of associated construction dust and debris within the subject structure.
5. As previously noted in is report, detectable levels of airborne particulates consisting primarily of skin cell fragments, pollen, and general debris particles were identified within the sampled interior areas surrounding the containment systems. Hence, and as an additional precautionary measure due to the presence of such materials, BioMax recommends that DGS considers the performance of supplemental post construction vacuuming and cleaning following the completion of interior renovation activities.
6. BioMax believes that any potential transmission and accumulation of the identified indoor airborne particulates may also be significantly reduced (if desired) on an immediate and ongoing basis through the implementation and use of routine HEPA filtered vacuuming and damp-wipe O&M cleaning methods employed by DGS maintenance personnel. BioMax's experience has indicated that these relatively simple measures and methods have been shown

to significantly reduce the accumulation of settled particulate debris on an immediate and ongoing basis if so desired.

7. Reasonable additional assessment and investigative measures may also be required upon the identification of new or previously undiscovered materials and/or information related to moisture/microbial impacts within the subject building structures, as necessary. Any occurrence and/or re-occurrence of moisture intrusion following routine O&M and/or general reconstruction within the subject building should also be reviewed and addressed through professional consultation, as necessary. BioMax is certainly prepared to provide such additional consultation pertaining to these and any follow-up investigative measures as necessary and upon request.

BioMax believes that the conclusions and recommendations outlined above are consistent with standard industry microbial mitigative and assessment practices as well as prudent industrial hygiene hazard control methods. Please do not hesitate to contact our offices directly at (510) 724-3100 if you have any additional questions, comments, or require further assistance regarding this matter.

Sincerely,



Michael A. Polkabla, CIH, REA
Vice President, Principal



LIMITATIONS

Please note that the professional opinions presented in this review are intended for the sole use of the California State Department of General Services (DGS) and their designated beneficiaries. No other party should rely on the information contained herein without the prior written consent of BioMax Environmental and DGS. The professional opinions provided herein are based on BioMax's review and understanding of current site information and observed site conditions present within the areas inspected at the time these services were performed. Professional recommendations provided as part of this limited scope of work are intended for client consideration only and are not intended as a professional or regulatory mandate. Implementation of any of the above measures or recommendations does not, in any way, warrant the day-to-day health and/or safety of building occupants, residents, site workers, nor regulatory or building code compliance status during normal and changing environmental conditions. As microbial contamination, by nature, may change over time due to additional moisture intrusion, favorable growth conditions, and changing environments, the findings of this report are subject to change in the event that such conditions and/or environments arise. Also, the professional opinions expressed here are subject to revision in the event that new or previously undiscovered information is obtained or uncovered.

The information contained in this and any other applicable communication is for consideration purposes only. It is not intended, nor should it be construed as providing legal advice or warranting any level of safety or regulatory compliance. The sole purpose of such information is to assist with the anticipation, identification, evaluation and control of elevated and/or unnecessary health of physical hazards. Any action taken based on this information, including but not limited to opinions, suggestions and recommendations, whether implied or expressed, is the sole responsibility of the individual taking the action. The management of acceptable health and safety is criteria dependent and situation specific in nature, therefore requiring extensive knowledge and prudent value assessments so as to be properly determined and maintained.

These services were performed by BioMax in accordance with generally accepted professional industrial hygiene principals, practices, and standards of care. Under the existing Industrial Hygiene Definition and Registration Act, all reports, opinions or official documents prepared by a Certified Industrial Hygienist (CIH) constitutes an expression of professional opinion regarding those facts or findings which are subject of a certification and does not constitute a warranty or guarantee, either expressed or implied.

**EMLab P&K**

Report for:

Mr. Michael Polkable
Blomax Environmental
775 San Pablo Ave.
Pinole, CA 94564

Regarding: Project: 032608-01; DGS, 450 N Street, Sacramento, CA
EML ID: 404691

Approved by:

A handwritten signature in black ink, appearing to read 'Dr. Kamashwaran Ramanathan'.

Lab Manager
Dr. Kamashwaran Ramanathan

Dates of Analysis:
Spore trap analysis: 03-31-2008

Project SOPs: Spore trap analysis (I100000)

This coversheet is included with your report in order to comply with AIHA and ISO accreditation requirements.

For clarity, we report the number of significant digits as calculated; but, due to the nature of this type of biological data, the number of significant digits that is used for interpretation should generally be one or two. All samples were received in acceptable condition unless noted in the Report Comments portion in the body of the report. Due to the nature of the analyses performed, field blank corrections of results is not a standard practice. The results relate only to the items tested.

EMLab P&K ("the Company") shall have no liability to the client or the client's customer with respect to decisions or recommendations made, actions taken or courses of conduct implemented by either the client or the client's customer as a result of or based upon the Test Results. In no event shall the Company be liable to the client with respect to the Test Results except for the Company's own willful misconduct or gross negligence nor shall the Company be liable for incidental or consequential damages or lost profits or revenues to the fullest extent such liability may be disclaimed by law, even if the Company has been advised of the possibility of such damages, lost profits or lost revenues. In no event shall the Company's liability with respect to the Test Results exceed the amount paid to the Company by the client therefor.

Document Number: 200091 - Revision Number: 5

EMLab P&K

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(650) 829-5800 Fax (650) 829-5852 www.emlab.com

Client: Biomax Environmental

C/O: Mr. Michael Polkaba

Re: 032608-01; DGS, 450 N Street, Sacramento, CA

Date of Sampling: 03-26-2008

Date of Receipt: 03-28-2008

Date of Report: 03-31-2008

SPORE TRAP REPORT: NON-VIABLE METHODOLOGY

Location:	13430732: Ambient at front entry		13430735: Ambient 12th floor balcony, SE		13430710: Ambient 23rd floor balcony, W		13430739: Hallway 24, outside cont 2424	
Comments (see below)	None		None		None		None	
Lab ID-Version†:	1777272-1		1777273-1		1777274-1		1777275-1	
	raw ct.	spores/m3	raw ct.	spores/m3	raw ct.	spores/m3	raw ct.	spores/m3
Alternaria								
Arthrinium								
Ascospores*	9	480	2	107	1	53		
Aureobasidium								
Basidiospores*	5	267	2	107				
Bipolaris/Drechslera group								
Botrytis								
Chaetomium					1	13		
Cladosporium	2	107			4	213		
Curvularia								
Epicoccum								
Fusarium								
Myrothecium								
Nigrospora								
Other colorless								
Penicillium/Aspergillus types†	8	427	2	107	2	107	2	107
Pithomyces								
Rusts*								
Smuts*, Periconia, Myxomycetes*			1	13				
Stachybotrys								
Stemphylium								
Torula								
Ulocladium								
Zygomycetes								
Background debris (1-4+)††	2+		2+		2+		3+	
Hyphal fragments/m3	< 13		13		< 13		< 13	
Pollen/m3	27		107		160		< 13	
Skin cells (1-4+)	< 1+		< 1+		< 1+		2+	
Sample volume (liters)	75		75		75		75	
TOTAL SPORE/m3		1,281		334		386		107

Comments:

* Most of these spore types are not seen with culturable methods (Andersen sampling), although some may appear as non-sporulating fungi. Most of the basidiospores are "mushroom" spores while the rusts and smuts are plant pathogens.

† The spores of *Aspergillus* and *Penicillium* (and others such as *Acremonium*, *Paecilomyces*) are small and round with very few distinguishing characteristics. They cannot be differentiated by non-viable sampling methods. Also, some species with very small spores are easily missed, and may be undercounted.

†† Background debris indicates the amount of non-biological particulate matter present on the trace (dust in the air) and the resulting visibility for the analyst. It is rated from 1+ (low) to 4+ (high). Counts from areas with 4+ background debris should be regarded as minimal counts and may be higher than reported. It is important to account for samples volumes when evaluating dust levels. The Limit of Detection is the product of a raw count of 1 and 100 divided by the percent read. The analytical sensitivity (counts/m3) is the product of the Limit of Detection and 1000 divided by the sample volume.

‡ A "Version" greater than 1 indicates amended data.

EMLab P&K

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Client: Biomax Environmental

C/O: Mr. Michael Polkabla

Re: 032608-01; DGS, 450 N Street, Sacramento, CA

Date of Sampling: 03-26-2008

Date of Receipt: 03-28-2008

Date of Report: 03-31-2008

SPORE TRAP REPORT: NON-VIABLE METHODOLOGY

Location:	13430694: Containment 2424, L		13430729: Containment 2424, R		13430705: Hallway containment 2402		13430701: 2402 containment, L	
Comments (see below)	None		None		None		None	
Lab ID-Version†:	1777276-1		1777277-1		1777278-1		1777279-1	
	raw ct.	spores/m3	raw ct.	spores/m3	raw ct.	spores/m3	raw ct.	spores/m3
Alternaria								
Arthrinium								
Ascospores*							1	53
Aureobasidium								
Basidiospores*								
Bipolaris/Drechslera group								
Botrytis								
Chaetomium								
Cladosporium								
Curvularia								
Epicoccum			1	13				
Fusarium								
Myrothecium								
Nigrospora								
Other colorless								
Penicillium/Aspergillus types†	2	107	1	53	1	53	1	53
Pithomyces								
Rusts*					1	13		
Smuts*, Periconia, Myxomycetes*								
Stachybotrys								
Stemphylium								
Torula								
Ulocladium								
Zygomycetes								
Background debris (1-4+)††	3+		3+		3+		3+	
Hyphal fragments/m3	< 13		< 13		< 13		< 13	
Pollen/m3	13		< 13		13		< 13	
Skin cells (1-4+)	2+		2+		3+		1+	
Sample volume (liters)	75		75		75		75	
TOTAL SPORE/m3		107		66		66		106

Comments:

* Most of these spore types are not seen with culturable methods (Andersen sampling), although some may appear as non-sporulating fungi. Most of the basidiospores are "mushroom" spores while the rusts and smuts are plant pathogens.

† The spores of *Aspergillus* and *Penicillium* (and others such as *Acremonium*, *Paecilomyces*) are small and round with very few distinguishing characteristics. They cannot be differentiated by non-viable sampling methods. Also, some species with very small spores are easily missed, and may be undercounted.

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The Limit of Detection is the product of a raw count of 1 and 100 divided by the percent read. The analytical sensitivity (counts/m3) is the product of the Limit of Detection and 1000 divided by the sample volume.

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Client: Biomax Environmental

C/O: Mr. Michael Polkable

Re: 032608-01; DGS, 450 N Street, Sacramento, CA

Date of Sampling: 03-26-2008

Date of Receipt: 03-28-2008

Date of Report: 03-31-2008

SPORE TRAP REPORT: NON-VIABLE METHODOLOGY

Location:	13430718: 2402 containmnet, R		13430706: Ambient 23rd floor, West balcony	
Comments (see below)	None		None	
Lab ID-Version†:	1777280-1		1777281-1	
	raw ct.	spores/m3	raw ct.	spores/m3
Alternaria				
Arthrinium				
Ascospores*				
Aureobasidium				
Basidiospores*			1	53
Bipolaris/Drechslera group				
Botrytis				
Chaetomium				
Cladosporium			2	107
Curvularia				
Epicoccum				
Fusarium				
Myrothecium				
Nigrospora				
Other colorless				
Penicillium/Aspergillus types†	2	107	1	53
Pithomyces				
Rusts*			1	13
Smuts*, Periconia, Myxomycetes*				
Stachybotrys				
Stemphylium				
Torula				
Ulocladium				
Zygomycetes				
Background debris (1-4+)††	3+		2+	
Hypheal fragments/m3	< 13		< 13	
Pollen/m3	< 13		107	
Skin cells (1-4+)	1+		< 1+	
Sample volume (liters)	75		75	
TOTAL SPORE/m3		107		226

Comments:

* Most of these spore types are not seen with culturable methods (Andersen sampling), although some may appear as non-sporulating fungi. Most of the basidiospores are "mushroom" spores while the rusts and smuts are plant pathogens.

† The spores of *Aspergillus* and *Penicillium* (and others such as *Acremonium*, *Paecilomyces*) are small and round with very few distinguishing characteristics. They cannot be differentiated by non-viable sampling methods. Also, some species with very small spores are easily missed, and may be undercounted.

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Client: Biomax Environmental

C/O: Mr. Michael Polkabila

Re: 032608-01; DGS, 450 N Street, Sacramento, CA

Date of Sampling: 03-26-2008

Date of Receipt: 03-28-2008

Date of Report: 03-31-2008

MoldRANGE™: Extended Outdoor Comparison**Outdoor Location: 13430732, Ambient at front entry**

Fungi Identified	Outdoor data	Typical Outdoor Data by Date†				Typical Outdoor Data by Location‡			
		Month: March				State: CA			
	spores/m3	low	med	high	freq %	low	med	high	freq %
Generally able to grow indoors*									
Alternaria	-	7	27	210	43	7	27	230	60
Bipolaris/Drechslera group	-	7	13	120	12	7	13	120	14
Chaetomium	-	7	13	120	8	7	13	110	19
Cladosporium	107	27	320	4,300	91	53	640	6,500	98
Curvularia	-	7	13	210	7	7	13	210	7
Nigrospora	-	7	13	110	7	7	13	170	8
Penicillium/Aspergillus types	427	27	160	1,600	82	40	210	2,500	88
Stachybotrys	-	7	13	310	3	7	13	330	5
Torula	-	7	13	170	8	7	13	150	13
Seldom found growing indoors**									
Ascospores	480	13	130	2,000	74	13	110	1,800	73
Basidiospores	267	13	320	5,700	90	13	270	6,900	95
Rusts	-	7	13	320	17	7	13	270	29
Smuts, Periconia, Myxomycetes	-	7	27	310	54	8	40	470	71
TOTAL SPORES/M3	1,281								

† The Typical Outdoor Data by Date represents the typical outdoor spore levels across North America for the month indicated. The last column represents the frequency of occurrence. The low, medium, and high values represent the 2.5, 50, and 97.5 percentile values of the spore type when it is detected. For example, if the frequency of occurrence is 63% and the low value is 53, it would mean that the given spore type is detected 63% of the time and, when detected, 2.5% of the time it is present in levels above the detection limit and below 53 spores/m3. These values are updated periodically, and if enough data is not available to make a statistically meaningful assessment, it is indicated with a dash.

‡ The Typical Outdoor Data by Location represents the typical outdoor spore levels for the region indicated for the entire year. As with the Typical Outdoor Data by Date, the four columns represent the frequency of occurrence and the typical low, medium, and high concentration values for the spore type indicated. These values are updated periodically, and if enough data is not available to make a statistically meaningful assessment, it is indicated with a dash.

*The spores in this category are generally capable of growing on wet building materials in addition to growing outdoors. Building related growth is dependent upon the fungal type, moisture level, type of material, and other factors. *Cladosporium* is one of the predominant spore types worldwide and is frequently present in high numbers. *Penicillium/Aspergillus* species colonize both outdoor and indoor wet surfaces rapidly and are very easily dispersed. Other genera are usually present in lesser numbers.

**These fungi are generally not found growing on wet building materials. For example, the rusts and smuts are obligate plant pathogens. However, in each group there are notable exceptions. For example, agents of wood decay are members of the basidiomycetes and high counts of a single morphological type of basidiospore on an inside sample should be considered significant.

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EMLab P&K

1150 Bayhill Drive, Suite 100, San Bruno, CA 94066
(650) 829-5800 Fax (650) 829-5852 www.emlab.com

Client: Biomax Environmental

C/O: Mr. Michael Polkabila

Re: 032608-01; DGS, 450 N Street, Sacramento, CA

Date of Sampling: 03-26-2008

Date of Receipt: 03-28-2008

Date of Report: 03-31-2008

MoldRANGE™: Extended Outdoor Comparison

Outdoor Location: 13430735, Ambient 12th floor balcony, SE

Fungi Identified	Outdoor data	Typical Outdoor Data by Date†				Typical Outdoor Data by Location‡			
		Month: March				State: CA			
	spores/m3	low	med	high	freq %	low	med	high	freq %
Generally able to grow indoors*									
Alternaria	-	7	27	210	43	7	27	230	60
Bipolaris/Drechslera group	-	7	13	120	12	7	13	120	14
Chaetomium	-	7	13	120	8	7	13	110	19
Cladosporium	-	27	320	4,300	91	53	640	6,500	98
Curvularia	-	7	13	210	7	7	13	210	7
Nigrospora	-	7	13	110	7	7	13	170	8
Penicillium/Aspergillus types	107	27	160	1,600	82	40	210	2,500	88
Stachybotrys	-	7	13	310	3	7	13	330	5
Torula	-	7	13	170	8	7	13	150	13
Seldom found growing indoors**									
Ascospores	107	13	130	2,000	74	13	110	1,800	73
Basidiospores	107	13	320	5,700	90	13	270	6,900	95
Rusts	-	7	13	320	17	7	13	270	29
Smuts, Periconia, Myxomycetes	13	7	27	310	54	8	40	470	71
TOTAL SPORES/M3	334								

† The Typical Outdoor Data by Date represents the typical outdoor spore levels across North America for the month indicated. The last column represents the frequency of occurrence. The low, medium, and high values represent the 2.5, 50, and 97.5 percentile values of the spore type when it is detected. For example, if the frequency of occurrence is 63% and the low value is 53, it would mean that the given spore type is detected 63% of the time and, when detected, 2.5% of the time it is present in levels above the detection limit and below 53 spores/m3. These values are updated periodically, and if enough data is not available to make a statistically meaningful assessment, it is indicated with a dash.

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Client: Biomax Environmental

C/O: Mr. Michael Polkabila

Re: 032608-01; DGS, 450 N Street, Sacramento, CA

Date of Sampling: 03-26-2008

Date of Receipt: 03-28-2008

Date of Report: 03-31-2008

MoldRANGE™: Extended Outdoor Comparison

Outdoor Location: 13430710, Ambient 23rd floor balcony, W

Fungi Identified	Outdoor data	Typical Outdoor Data by Date†				Typical Outdoor Data by Location‡			
		Month: March				State: CA			
	spores/m3	low	med	high	freq %	low	med	high	freq %
Generally able to grow indoors*									
Alternaria	-	7	27	210	43	7	27	230	60
Bipolaris/Drechslera group	-	7	13	120	12	7	13	120	14
Chaetomium	13	7	13	120	8	7	13	110	19
Cladosporium	213	27	320	4,300	91	53	640	6,500	98
Curvularia	-	7	13	210	7	7	13	210	7
Nigrospora	-	7	13	110	7	7	13	170	8
Penicillium/Aspergillus types	107	27	160	1,600	82	40	210	2,500	88
Stachybotrys	-	7	13	310	3	7	13	330	5
Torula	-	7	13	170	8	7	13	150	13
Seldom found growing indoors**									
Ascospores	53	13	130	2,000	74	13	110	1,800	73
Basidiospores	-	13	320	5,700	90	13	270	6,900	95
Rusts	-	7	13	320	17	7	13	270	29
Smuts, Periconia, Myxomycetes	-	7	27	310	54	8	40	470	71
TOTAL SPORES/M3	386								

† The Typical Outdoor Data by Date represents the typical outdoor spore levels across North America for the month indicated. The last column represents the frequency of occurrence. The low, medium, and high values represent the 2.5, 50, and 97.5 percentile values of the spore type when it is detected. For example, if the frequency of occurrence is 63% and the low value is 53, it would mean that the given spore type is detected 63% of the time and, when detected, 2.5% of the time it is present in levels above the detection limit and below 53 spores/m3. These values are updated periodically, and if enough data is not available to make a statistically meaningful assessment, it is indicated with a dash.

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Client: Biomax Environmental

C/O: Mr. Michael Polkabila

Re: 032608-01; DGS, 450 N Street, Sacramento, CA

Date of Sampling: 03-26-2008

Date of Receipt: 03-28-2008

Date of Report: 03-31-2008

MoldRANGE™: Extended Outdoor Comparison

Outdoor Location: 13430706, Ambient 23rd floor, West balcony

Fungi Identified	Outdoor data	Typical Outdoor Data by Date†				Typical Outdoor Data by Location‡			
		Month: March				State: CA			
	spores/m3	low	med	high	freq %	low	med	high	freq %
Generally able to grow indoors*									
Alternaria	-	7	27	210	43	7	27	230	60
Bipolaris/Drechslera group	-	7	13	120	12	7	13	120	14
Chaetomium	-	7	13	120	8	7	13	110	19
Cladosporium	107	27	320	4,300	91	53	640	6,500	98
Curvularia	-	7	13	210	7	7	13	210	7
Nigrospora	-	7	13	110	7	7	13	170	8
Penicillium/Aspergillus types	53	27	160	1,600	82	40	210	2,500	88
Stachybotrys	-	7	13	310	3	7	13	330	5
Torula	-	7	13	170	8	7	13	150	13
Seldom found growing indoors**									
Ascospores	-	13	130	2,000	74	13	110	1,800	73
Basidiospores	53	13	320	5,700	90	13	270	6,900	95
Rusts	13	7	13	320	17	7	13	270	29
Smuts, Periconia, Myxomycetes	-	7	27	310	54	8	40	470	71
TOTAL SPORES/M3	226								

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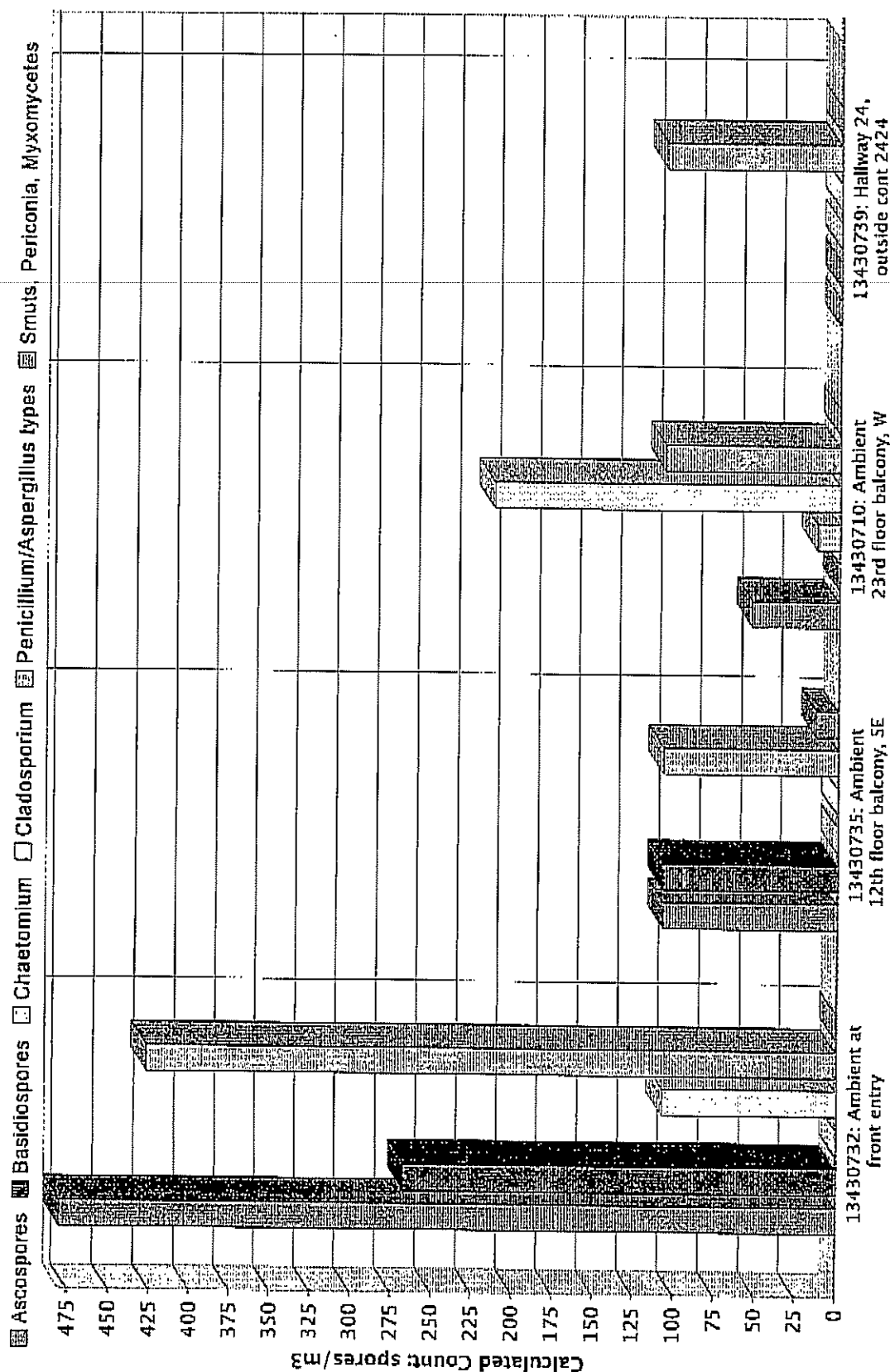
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03-31-2008: 032608-01

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SPORE TRAP REPORT: NON-VIABLE METHODOLOGY



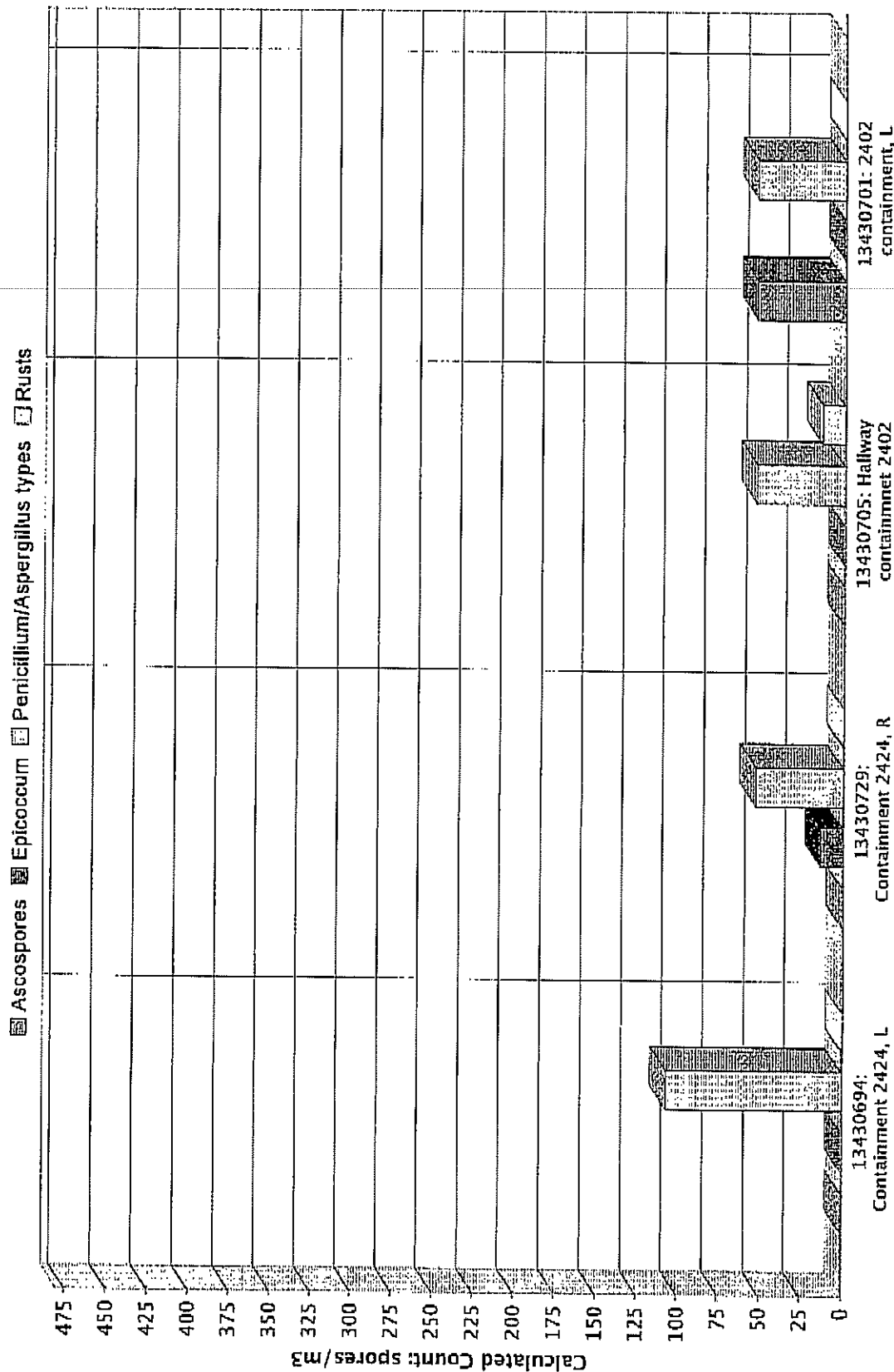
Comments:

Note: Graphical output may understate the importance of certain "marker" genera.

03-31-2008: 032608-01

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SPORE TRAP REPORT: NON-VIABLE METHODOLOGY



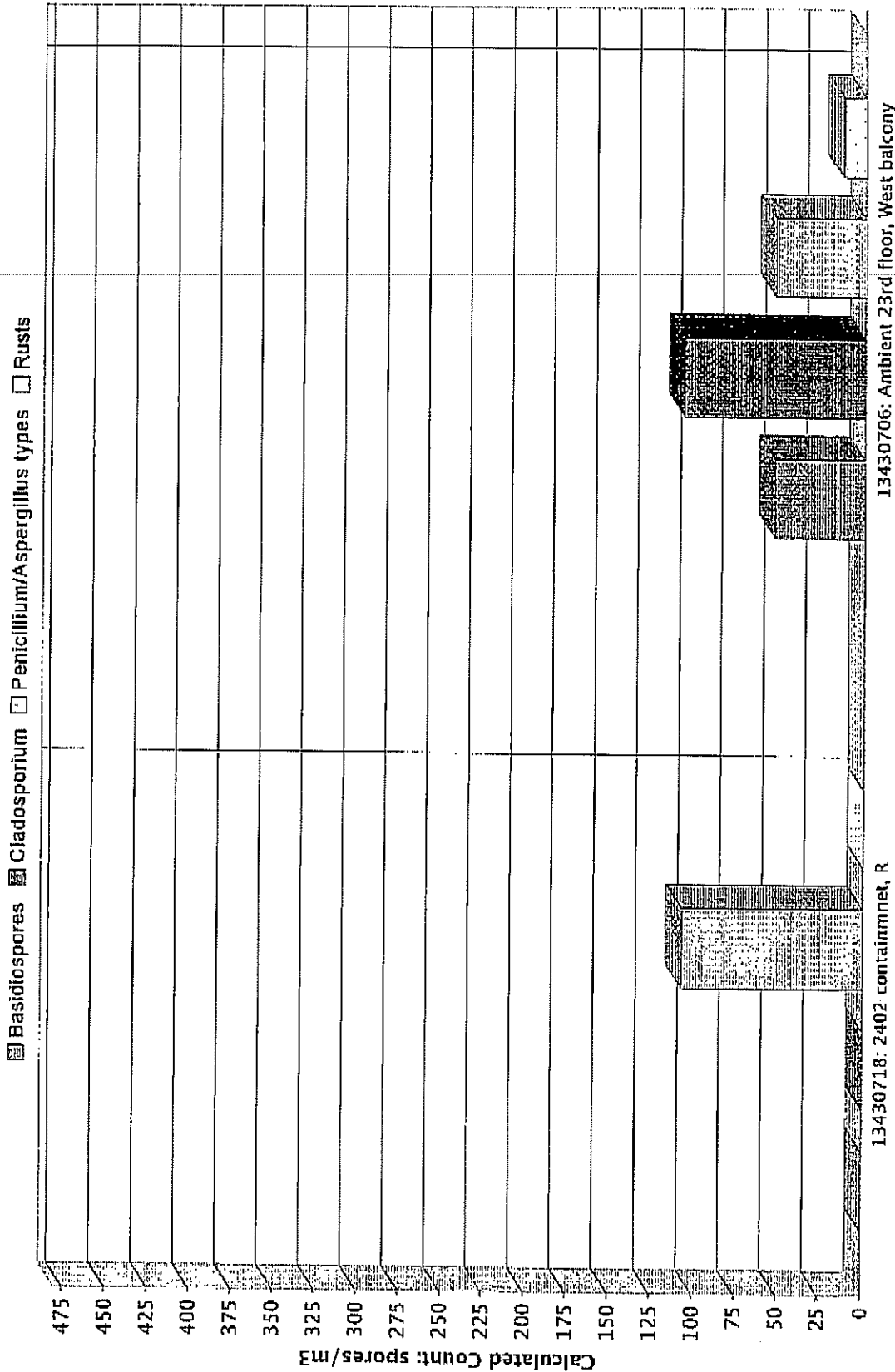
Comments:

Note: Graphical output may understate the importance of certain "marker" genera.

03-31-2008: 032608-01

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SPORE TRAP REPORT: NON-VIABLE METHODOLOGY



Comments:
Note: Graphical output may understate the importance of certain "marker" genera.

MICROBIAL SPORE TRAP AIR SAMPLING RECORD

Page 1 of 1

BioMax Environmental
775 San Pablo Ave.
Pinole, CA 94564

www.biomaxenvironmental.com

Phone: (510) 724-3100
Fax: (510) 724-3145
biomaxenv@aol.com

Location: 450 N-Street Sacramento, CA	Client: DGS Project #: 032608-01
Date: 3/26/08	Laboratory: EMCLabs
Collected by: MA Palkala	Req. Turn Around: 24 hr
Signature: <i>MA Palkala</i>	Analysis (circle): <u>Fungal</u> Particulate <u>ID / Quantification.</u>

Sample Number	Time	Location/Desc.	Temp / RH
13430732	0930	Ambient @ Front Entry	46°F / 55.5%
13430735	0950	Ambient 12th Floor Balcony (SE)	61°F / 37%
13430710	1010	Ambient 23rd Floor Balcony (W)	54°F / 53%
13430739	1525	Hallway 24 Outside Cont ²⁴²⁴	74°F / 27% RH
13430694	1538	Contaminant 2424 L	75 / 28 %
13430729	1545	Contaminant 2424 R	75 / 28 %
13430705	1605	Hallway Contaminant 2402	77 / 25 %
13430701	1610	2402 Contaminant L	78 / 25 %
13430718	1615	2402 Contaminant R	77 / 25 %
13430706	1630	Ambient 23rd Floor West Balcony	66 34%
Total Sample Time (min): 5	Flow Rate (l/min): 15	Total Sample Volume (liters): 75L	Ambient Conditions: Clear / Mild 0-5 NW Comments:

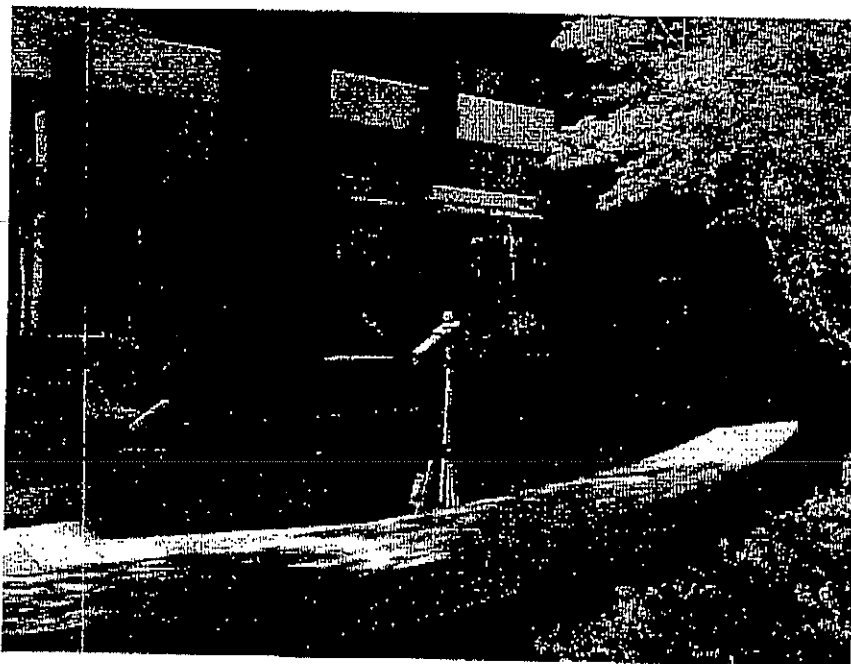
Please sign this form below acknowledging sample receipt and return executed form with laboratory reports. Fax, send, e-mail results to BioMax Environmental at (510) 724-3145 biomaxenv@aol.com
Other Instructions:

Relinquished by: <i>MA Palkala</i>	Received By:
Method of Transportation: Fed X	
Time/Date Sent: 3:00 3/27/08	Time/Date Received:

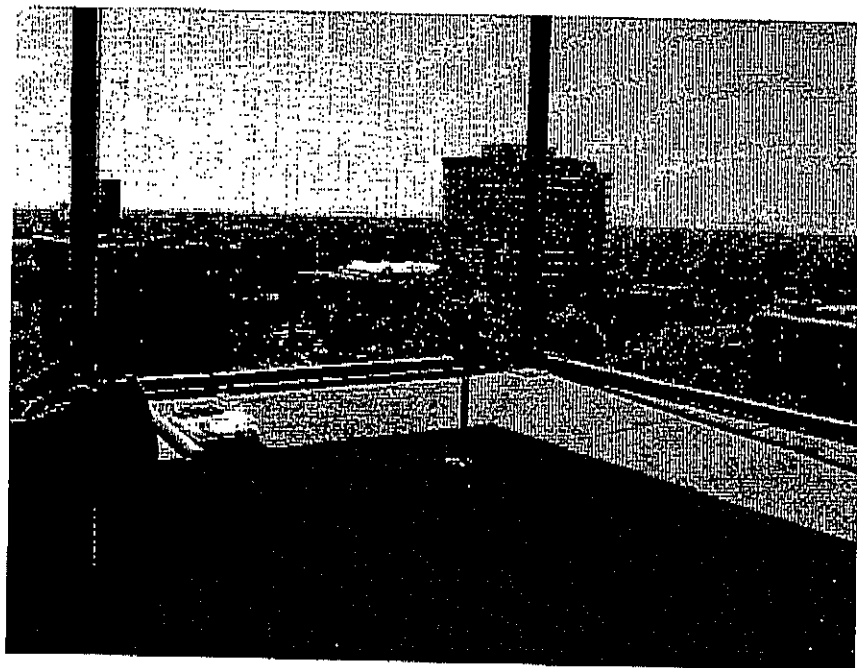
Attachment A: Digital ImagesMarch 26th, 2008BOE Building 24th Floor Break Rooms

Sacramento, CA

Page 1 of 5

[Click here for color photos](#)

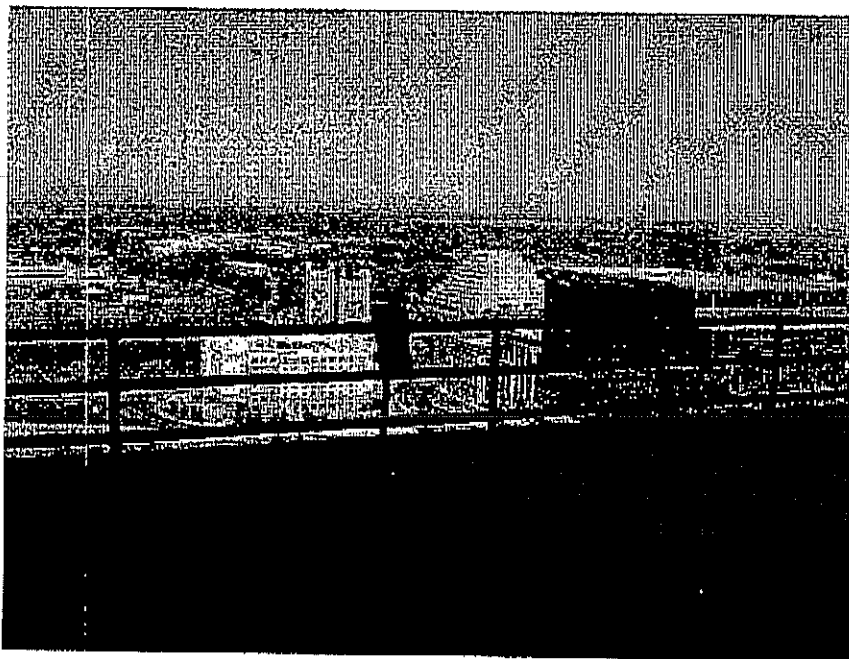
- 1) Image of ambient air sampling location at front entry of BOE Building (Subject Building) located at 450 N Street, Sacramento, California at time of assessment.



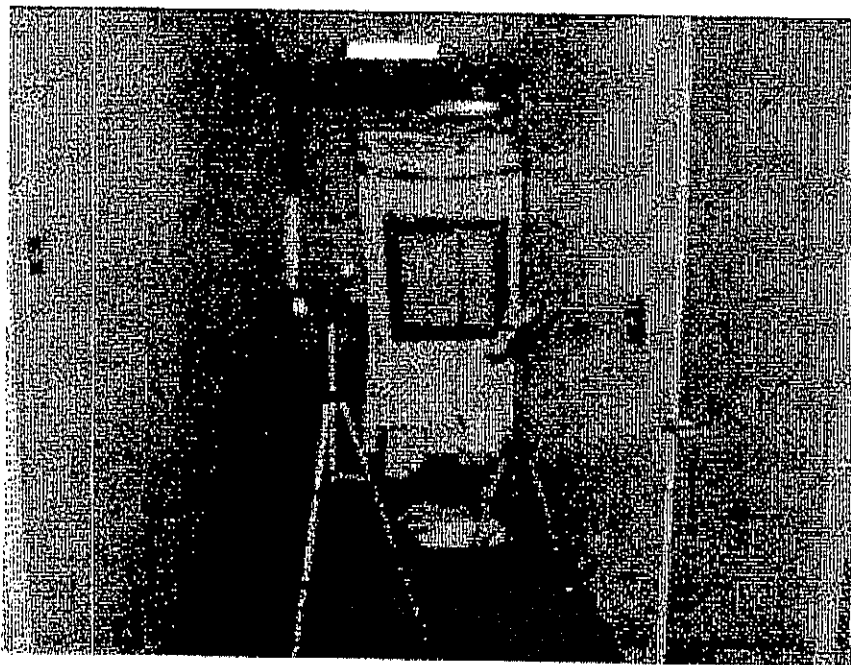
- 2) Image of ambient air sampling location on 12th Floor southeast deck area at time of assessment.

March 26th, 2008
BOE Building 24th Floor Break Rooms
Sacramento, CA

Page 2 of 5



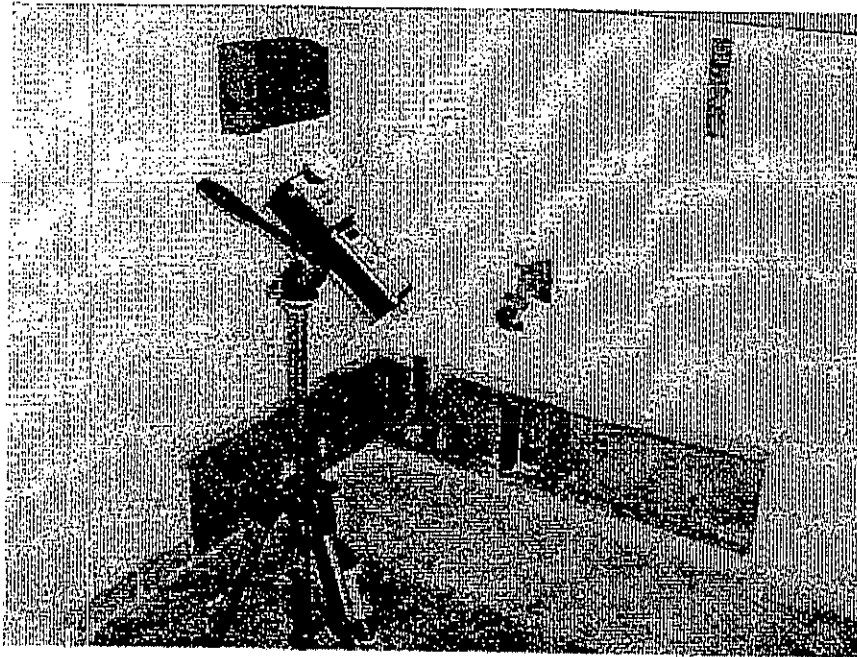
3) Image of ambient air sampling location on 23rd floor western deck area at time of assessment.



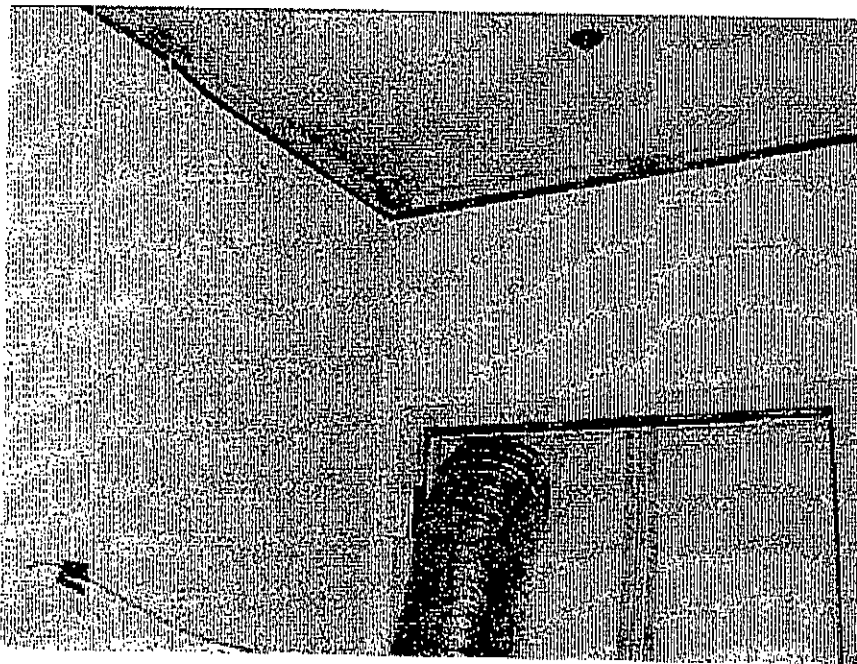
4) Image of hallway air sampling location adjacent to Room 2424 containment (outside containment) at time of assessment.

March 26th, 2008
BOE Building 24th Floor Break Rooms
Sacramento, CA

Page 3 of 5



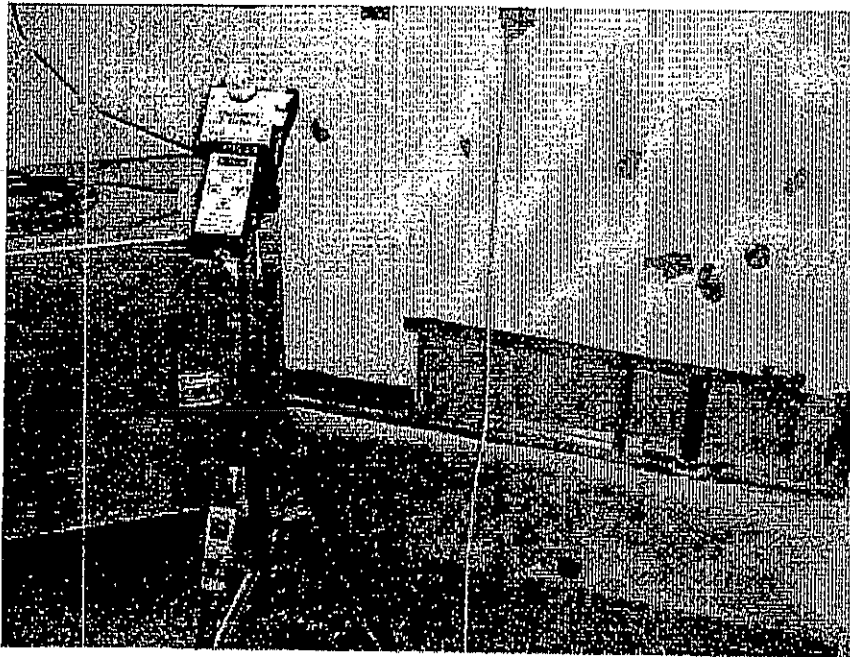
- 5) Image within 2424 containment indicating location of air sampling equipment and removal of cabinetry and wallboard materials.



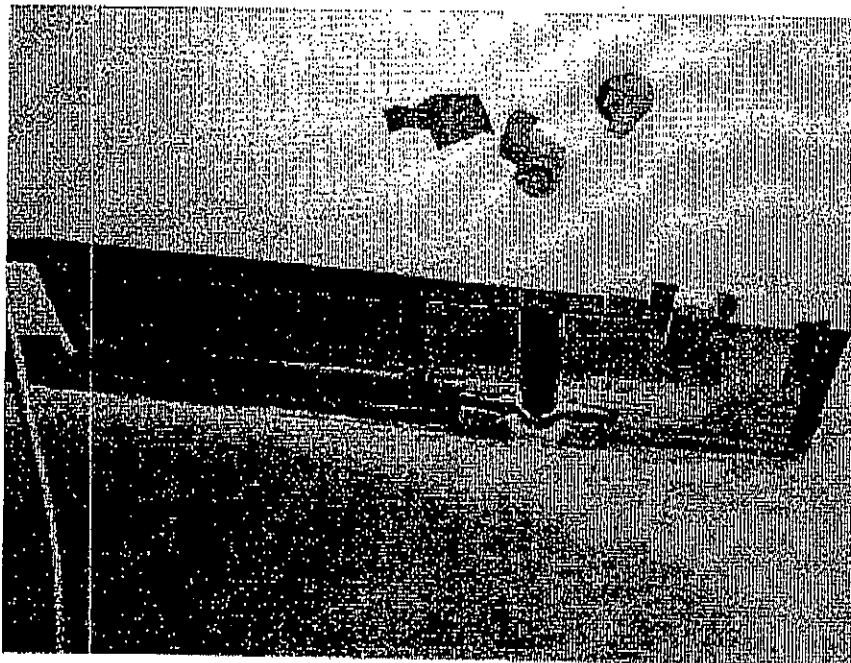
- 6) Image within 2424 containment viewing ceiling containment and exhaust ventilation at entry/exit chamber doorway.

March 26th, 2008
BOE Building 24th Floor Break Rooms
Sacramento, CA

Page 4 of 5



- 7) Image within 2402 Break Room containment indicating location of air sampling equipment and physical cabinet and wallboard removal at time of clearance assessment.



- 8) Close-up image of location within 2402 containment where cabinetry and wallboard materials were removed by mitigation contractor.

March 26th, 2008
BOE Building 24th Floor Break Rooms
Sacramento, CA

Page 5 of 5



9) Image of Hygientech (HTI) technician collecting surface tape sample at time of assessment within Break Room 2402 containment.



10) Image of Hygientech (HTI) technician collecting parallel spore trap air sample on 23rd Floor West Balcony following interior assessment.